(12) United States Patent Chino et al.

(10) Patent No.:

US 6,403,720 B1

(45) Date of Patent:

Jun. 11, 2002

(54)	RUBBER COMPOSITION AND
,	PRODUCTION PROCESS THEREOF

- (75) Inventors: Keisuke Chino; Hidekazu Onoi; Satoshi Mihara, all of Hiratsuka (JP)
- Assignee: The Yokohama Rubber Co., Ltd., Tokyo (JF)
- Subject to any disclaimer, the term of this (*) Notice: patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.
- (21) Appl. No.: 09/642,635

Aug. 31, 1999

(22)Filed: Aug. 22, 2000

Foreign Application Priority Data (30)

) (IP) maintenation in the comment of the comment o	023750
L7 C08L 7/06; C08	
C08L 23/00; C08I	

(JP) 11-246198

525/219, 232, 233, 238, 240, 241

U.S. CL 525/191; 525/213; 525/219; (52)525/232; 525/233; 525/238; 525/240; 525/241 Field of Search 525/191, 213.

(56)

30

References Cited

U.S. PATENT DOCUMENTS

4.146.592 A *	3/1979	Kitagawa et al 525/213
5,395,891 A *	3/1995	Obrecht et al 525/194
5,184,295 B1 *	2/2001	Obrecht at at
6,242,534 Bi *	5/2001	Obrecht et al 525/191

FOREIGN PATENT DOCUMENTS

10204217 A 8/1998

* cited by examinar

Primary Examiner-Nathan M. Nutter (74) Attorney, Agent, or Firm-Arent, Fox, Kintney, Plotkin & Kabu

ABSTRACT

A rubber composition containing 50 to 90 parts by weight of a diseas rubber and 50 to 10 parts by weight of a gelled rubber having a toluene swelling index of 16 to 150, based upon the total amount of 100 parts by weight of the diene rubber and the gelled rubber.

4 Claims, No Drawings

5

3

RUBBER COMPOSITION AND PRODUCTION PROCESS THEREOF

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention relates to a subber composition suitable for use as, for example, a tire, which is obtained by blending a gelied rubber symbosized with an acid anhydride esc., without substantially decreasing abrasion resistance, rolling resistance, and a process for producing the same.

2. Description of the Related Art

In rubber compositions used for automobile tires etc., rubber compositions superior in drivability on wet road and 15 low in miling resistance are desired. From such a viewpoint, for example, Japanese Unexamined Patent Publication (Kokai) No. 10-204217 proposes to blood an SBR rubber gel to a nubber composition. This SBR rubber gel is synthesized by a method of cross-linking the SBR rubber with a poly- 20 functional compound such as divinylbenzene during the polymerization or a method of cross-linking of the polymer after polymerization with a peroxide etc.

SUMMARY OF INVENITION

It is known that, when carbon is blended into rubber the tan & curve with respect to the temperature becomes broad and the viscoelastic properties are deteriorated. The present inventors engaged in research to develop a rubber composition having a superior wet performance, that is, having a 30 high grip on was road surfaces and having a low rolling resistance, by mixing a rubber having a low glass transition temperature Tg with a gelled subber having a high glass transition temperature Tg so as to suppress the incorporation of carbon black to high Ig nibber and hopefully to obtain a 35 rubber composition with the balanced values of tan 5 at 0° C. and tan 8 at 60° C. As a result, we found that, by mixing a galled subber obtained by gelation with a cross-linking agent in the diene rubber, the viscoelastic properties of the rubber composition can be improved. Further, we found that 40 a public composition prepared by a two-stage mixing method of premixing the carbon black and other compounding agents to a low Tg ribber, and then mixing the resultant mixture with the gelled subber obtained by gelstion of a high To rubber further provides the improved viscoelastic prop- 45 erries

Accordingly, the objects of the present invention is to provide a rubber composition having a superior tan 8 balance, without substantially decreasing the abrasion resistance and with the improved wei performance and decreased 50 rolling resistance and a process for producing the same.

In accordance with the present invention, there is provided a rabber composition comprising 50 to 90 parts by weight of a diene rubber and 50 to 10 parts by weight of a galled rubber having a toluene swalling index of 16 to 150, ss based upon the total amount of 100 parts by weight of the diene nibber and the gelled rubber.

In accordance with the present invention, there is also provided a process for producing a rubber composition comprising 100 parts by weight of a starting rubber con- 60 taining 50 to 90 parts by weight of a diene mibber and 50 to 10 parts by weight of a gelled rubber, the glass transition temperature of the diene rubber being at least 10° C. less than the transition temperature of the gelled rubber, and 30 to 120 parts by weight of a filler, comprising the steps of es mixing the diene rubber and at least 80% by weight of the total amount of the filler at a imperature of at least 135° C.,

and then mixing the gelled rubber and the remaining filler with the resultant mixture.

DESCRIPTION OF THE PREFERRED EMBODIMENT

In a rire use rubber composition comprised of a low Tg (i.e. glass transition temperature) polymer and high Tg polymer, the tan 8 around 0° C. is affected to a large extent by the compatibility of the two polymers. The tan 6 around with the improved wet performance, and with the dacreased 10 0° C. is improved if the two polymers are incompatible. Further, by blending carbon black to the rubber, the tan o curve is broadened. Therefore, whon the carbon black is predominated in the low Tg polymer and then the high Tg polymer is blended thereto, the incorporation of the carbon black into the high Tg polymer side is suppressed and the tau 8 around 80° C, is decreased.

In the present invention, a gelled rubber having a toluene swelling index of 16 to 150 is blended, as the high Tg polymer, to a low Tg polymer of a diene subber the incompstibility of the two polymers is promoted, the tan 8 around 0° C. is improved, and the abrasion resistance is not too much impaired or, in some cases, is even improved. Further, according to the present invention, when the filler such as the carbon black is predominated in the low Tg polymer and when the gelled rubber is added thereto, the tan & around 0° C. is improved and the tan 8 around 60° C, is decreased and the abrasion resistance is maintained.

The gelled rubber used in the present invention may be produced by, for example, reacting a diene rubber such as a styrene-butadiene copolymer rubber (SBR) with maleic anhydride and a phenol compound having an aromatic hydroxyl group (e.g., phenoi, t-butylpheno), catechol and t-butyl catechol, hydroquinone, and resorcinel etc.) or a perexide compound (e.g., dicumyl perexide etc.) or a mercapto compound (e.g., inthiocyanic acid, 1,10-decanethio), etc.) at a temperature of 80 to 250° C.

Examples of the above phenol compounds are as follows. Monophenol Compounds

2,6-di-t-butyl-p-cresol butylated hydroxyanisole (BHA)

2.6-di-t-butyl-4-ethylphenol

Stearyl-8-(3,5-di-t-butyl-4-hydroxyphecyl) propionate Bisphenol Compounds

2,2'-methylenebis(4-methyl-6-t-butylphenol)

2.2'-methylenebis(4-ethyl-6-t-butylphenol)

4,4'-thiobis(3-methyl-6-t-butylphenol)

4,4'-butylidenebis(3-methyl-6-t-butylphenol)

3,9-bis[1,1-dimethyl-2-[8-(3-t-butyl-4-bydroxy-5methylphenyl)propionyloxy]ethyl]2,4,8,10tetraoxapyro[5,5]undecane

Phenol Polymer Compounds

1,1,3-mis(2-methyl-4-hydroxy-5-t-butylphenyl)butane

1,3,5-trimetby1-2,4,6-tris (3,5-di-t-buty1-4hydroxybenzyl)benzene

Tetrakis-(methylene-3-(3',5'-di-t-butyl-4-bydroxyphenyi) propiousts Junethane

Bis[3,3'-bis(4'-bydroxy-3'-t-butylphenyl)butyric acid] givool ester

1,3,5-tris(3 ', 5'-di-t-buty)-4'-bydroxybenzyl)-S-triazine-2,4,6-(1H,3H,5H) trione

Tocopherol(S)

Other Phenoi Compounds

2,6-di-terr-butyl-4-methylphenol (BHT)

Mono(or di-or tri-)(o-methylbenzyl)phenoi

2,2'-methylenebis(4-methyl-6-tert-butylphenol)

3

2,2-mothylonobis(4-ethyl-6-tert-butylphenol) 4,4-butylidensbis(6-tert-butyl-3-methylphenol)

4,4'-thiobis(6-tert-butyl-3-methylphenol)

1,1-bis(4-hydroxyphenyl)-cyclohexane

2,5-di-tert-butylbydroquinone

2,5-di-terr-smylhydrogulgone

comprises the phenol compound abstraction of the hydrogen from the benzyl position or allyl position, adding the benzyl radicals or allyl radicals thus formed to the acid anhydride, and adding the radicals to the double bonds of other rubber molecules or radical coupling them with other rubber molecules to form cross-linking.

The gelled subber according to the present invention has to have a toluene swelling index of 16 to 150, preferably 16 to 100. The "tolinene swelling index" in the present invention 15 is determined by immersing 0.1 g of a gelled rubber in 100 mi of toluene at room temperature for 24 hours, weighing the weight of the rubber at that time (wet weight), then drying the rubber in vacuo at room temperature for 24 hours, measuring the dry weight, and finding the swelling index 20 from the wes weight/dry weight. If the toluene swelling index is too small, the abrasion resistance deteriorates, whereas if it is conversely too large, there is no great difference with normally ungelled subper and no improvement effects in the viscoelastic properties are observed. Further, a gelled rubber containing at least 0.1% by weight, 25 preferably 0.5 to 10% by weight, of an acid anhydride molety in the molecule, has less deterioration of the abrasion resistance and in some cases is further improved compared with an ordinary subber. Further, a rabber composition obtained by mixing the gelled rubbet produced with a 30 peroxide compound or mercapto compound to a dione rubber may be affected in vulcanized physical properties with the peroxide compound or mercapto compound, and therefore, a gelled subber produced from an acid subydride and phenol compound is more preferable.

The rubber composition according to the present invention is obtained by blending 50 to 90 parts by weight, preferably 60 to 85 parts by weight, of the diene rubber and 30 to 10 parts by weight, preferably 40 to 15 parts by weight, of at least one gelled rubber, (Note: the total amount of 100 parts by weight). If the amount

Examples of the other cross-linking agents are organic peroxides such as dicumyl peroxide, t-butylcumyl peroxide, bis-(t-butyl-peroxy-isopropyl)henzene, di-t-butyl peroxide, 2,5-dimethylbexane-2,5-dihydroperoxide, 2,5-dimethyl-3bexise-2,5-dibydroperoxide, benzoyl peroxide, 2,4- 45 dichlorobenznyl peroxide, t-butyl perbenzoate; organic azo compounds such as azobisisobutyronitrile and azobiscyclohexanenitrile; dimercapto compounds and polymercapto compounds such as dimercapteethane, 1,6dimercaptohexane, and 1,3,5-trimercaptotriazine, trimethy- so iol propane tris(β-thiopropionate), 1,8-dimercapto-3,6dioxacctane, mercapto-terminated polysulfide rubbers such as mercapio-terminated reaction products of bischloroethyliormal and sodium polysulfide, sulfur chloride, dimercapto scids, quinones (e.g., p-quinone), quino- 55 nedioximes (e.g., p-quinonedioxime, p-quinoneoxime beazoate), polyhalides (trichloromelamine, hexachiorocyclopentadiene, octachiorocyclopentadiene, trichloromethanesulfochloride, benzotrichloride, paraffin chloride, FVC, chloroprene rubber, chlorosulfunated so polyethylene, etc.), and metal oxides (stantous chloride, lead oxide, etc.), or triethanolamine, paraformaldehyde or polyoxymethylene and protonic acid or Lewis acid (stannic (II) chloride, paratoluene sulfonic acid, etc.), boranes (for example, niethylamine-chloroborane, methylene diamine- 65 bischloroborane); dinitrones (phonylhydroxylamine or terephthalaldehyde), diminile oxides (terephthalonitrile

oxide, terephthalehydroxamyl chloride), dininile imines (terephthalylphenyllydrazide chloride and triethylamine), disydnones (p-phenylene-3,3'-disydnones), thionyl paraphenylene diamine, etc.

In the present invention, at least 0.1% by weight, preferably 0.5 to 30% by weight, of the maleic anhydride and at least 0.05% by weight, preferably 0.1 to 5% by weight, of a phenol compound such as test-butyl catechol are reacted with the rubber. The reaction mechanism of the gelled rubber blended is too small, the expected effect is hard to appear, while if conversely too large, the abrasion resistance is deteriorated.

The diene rubber used in the rubber composition of the present invention is not particularly limited, but any diene rubber generally used in various types of rubber compositions in the past, such as natural rubber (NR), polyisoprene rubber (IR), styrene-butadiene copolymer rubber (SHR), polybutadiene rubber (BR), scrylonifile butadiene copolymer rubber (NBR), butyl rubber (IIR), chloroprene rubber, ethylene-propylene chese copolymer rubber, ethylene-propylene diene copolymer rubber, etc. may be mentioned. These rubbers may be used alone or in any blend thereof.

The rubber composition according to the present invention is preferably produced as follows.

That is, according to the present invention, 30 to 120 parts by weight, preferably 40 to 100 parts by weight, of a filler such as carbon black or silica is blended to 100 parts by weight of a starting nubber comprised of 50 to 90 parts by weight, preferably 50 to 85 parts by weight, of a diene rubber and 50 to 10 parts by weight, preferably 40 to 15 parts by weight, of a gelled rubber. At that time, a diene rubber having a glass transition temperature Tg of at least 10° C lower than the Tg of the gelled rubber is preferable. More preferably, the diene rubber and at least 80% by weight of the total weight of the filler are mixed at a temperature of at least 135° C., and then the gelled rubber and the remaining filler are mixed to produce the desired rubber composition.

The subber composition may contain thereof compounding agents normally used in the subber industry if necessary. As such compounding agents, in addition to a filler such as the above carbon black and silica, for example a vulcanization accelerator, vulcanization promoter, astioxidant, plasticizer, softener, etc. may be mentioned. These may be blended in their respectively necessary amounts.

The rubber composition according to the present invention may be used for various types of rubber products such as tires, hoses, conveyor belts, rubber sheets, and fenders, but is particularly preferably used as a rubber composition for a tire.

EXAMPLES

The present invention will now be further illustrated by, but is by no means limited to, the following Examples.

> Standard Example 1, Examples 1 to 2, and Comparative Example 1

Synthesis of Gelled Rubber

(1) SBR1-gel 1: Gelled rubber SBR1-gel 1 was produced by adding, to 500 g of styrene-butadiene copolymer rubber (SBR) (glass transition temperature: ~24.5° C.) (NS110 manufactured by Nippon Zeou), 5 g of malcic anhydride and 0.5 g of tert-butyl catechol, followed by mixing in a kneader at 150° C. for about 60 minutes. When 0.2 g of the rubber thus obtained was immersed in 100 mi of toluene for 24 hours, the rubber was swelled, without dissolving, and therefore the gelation was confirmed. The swelling index was calculated by the wet weight/dry weight. Further, from IR analysis, absorption at 1760 cm⁻¹ was confirmed, whereby the incorporation of an acid subydride structure to the rubber was confirmed.

-5

The SBR1-gel 1 thus obtained had an acid anhydride content of 0.8% by weight and a swelling index of 18.

(2) SBR1-PO gel: 250 g of SBR (NS110 manufactured by Nippon Zeon) was dissolved in 1 liter of cyclohexane. 20 g of dictional peroxide was added thereto. The mixture was 5 stirred in an autoclave under mitrogen at 60° C. for 2 hours, then the temperature was raised and the mixture etirred at 150° C. for 45 minutes. Next, 100 g of a 10% aqueous sulfuric acid solution was added to congulate the gelled rubber and the resultant product was dried in vacuo at 60° C. 10 over 2 days. The swelling index was 13.

(3) SBR2-gel 1: Produced from styrene buradiene rubber (NS110, Nippon Zeon) in a similar method to the production of SBR1-gel 1 except for mixing at 150° C, for 30 minutes. The swelling index was 18.

The gelied rubber synthesized above and the starting rubber SBR1 (NS110 manufactured by Nippon Zenn, Tg =-24.5° C.) were used to obtain, by mixing, the rubber compositions of Standard Example 1, Comparative Example 1, and Examples 1 to 2 by a Banbury mixer and a roll mill. 20 based upon the formulations (parts by weight) shown in Table I and ordinary methods. The rubber compositions thus obtained were press vulcanized at 150° C. for 20 minutes to prepare the desired test pieces which were then evaluated for physical properties.

The ingredients of the formulations used were as follows: NR (natural rubber): glass transition temperature of -56°

Carbon black: Seast 3H (Tokai Carbon)

Zinc white: Zinc White No. 3 (Seido Chemical)

Steame acid: Lunae YA (Kao Soap)

Amioxidant 6C: Noctac 6C (Ouchi Shinko Chemical) Sulfur: Oil extended sulfur (Karuizawa Refinery)

Accelerator: Vulcanization accelerator Noccelar CZ 35 (Ouchi Shinko Chemical)

The physical properties were evaluated and tested by the following methods. The results are shown in Table I.

tan 8 (0° C., and 60° C.): A spectrometer (manufactured by Toyo Seiki Seisakusho) was used for measurement at an amplitude of e2%, a vibration of 20 Hz, and an initial strain of 10%. The larger the value of the tan 8 (0° C.), the higher the grip on wet road surfaces exhibited, while the smaller the value of the tan 8 (60° C.), the smaller the rolling resistance with ideal.

Abrasion resistance: A Lambourn abrasion tester manufactured by Iwamoto Seisakusho was used for measurement according to IIS K6264 and the value shown indexed to the value of Comparative Example 1 as 100. The larger the value, the better the abrasion resistance.

TABLE I

	Standard Ex. 1	Comp. Ex. 1	Exc. 1	Ex. 2					
NB:	ं रह	70-	70	70					
SBR1302	30	***	~~						
SBRIAPO gel		30		****					
SBR1-gel 1			30						
SBR2-get 1			***	30 50					
Cuthou black	-50	20	30	50					
Zioc white	ä	3	3	3					
Stearic acid	2	3	3	j.					
Actionidani (C)	2	3	1	1					
Sultur	1,78	1.75	1.75	1.75					
Vulnamization accelerator	3	7.	3.	1					
හැර (0° C.)	0.282	0.361	0.305	0.334					

6

TABLE 1-continued

	Stoudard Ex. 1	Comp. Ex. 1	Ex. I	Ex. 3
:mð (60° C.)	0.176	0.788	0.165	0,138
tand balance	3,602	1.603	1.848	1,987
Abrasion resistance (index)	100	§ 4	94	97

Standard Example 2, Examples 3 to 5, and Comparative Examples 2 to 3

Rubber compositions having the formulations (parts by weight) shown in Table II were obtained by similar methods to Example 1 and Comparative Examples 1 to 2, except for using the following SBR3 and 4 as the diene rubber and the following rubbers as the gelled rubber and evaluated the physical properties thereof. The results are shown in Table II.

SBR3: Styrene butadieno rubber (Asahi Chemical, Tufden 1000K, Tg. -72° C.)

SBR4: Styrene butadiene rubbet (Nippon Zeon, Nipol 9529, Tg: -21° C.) SBR4-PO gel 1: Prepared by an analogous method to SBR1-PO gel by dissolving 250 g of styrene-butadiene rubbet (Nippon Zeon, Nipol 9550) in 1 liter of cyclohexane and adding 37.5 g of dicumyl peroxide. The swelling index was 13.

SBR4-gel 1: A gelled rubber prepared from styrene-butadiene rubber (Nippon Zeon, Nipol 9550) in an analogous method to SBR1-gel 1. The swelling index was 14

SBR4-PO gel 2: Prepared by an analogous method to SBR1-PO gel by dissolving 250 g of styrene-butadiene rubber (Nippon Zeon, Nipol 9550) in 1 liter of cyclohexane and adding 20 g of dicumyl peroxide. The swelling index was 18.

SBR4-gel 2: A gelled subber prepared from styrene-butadiene subber (Nippon Zeon, Nipol 9550) in an analogous method to SBR2-gel. The swelking index was 25.

SBR5-gel: Styrene-butadiene rubber gel (Nippon Zeon, 45, 2001). The swelling index was 72.

TABLE II

					*********	-
	Standard E. 23	Comp. Ex. 2	Comp. Ex. 3	Ex. 3	Ex. 4	Ex. 9
SBR3	70	70	70	70	70	70
SBR4	90	•		****	agent :	
SBR4-PQ gel 1	and the second	30				••••
SBR4-gel 1.		·	30	···		,
SBR4-PO gel 2				30	*****	~ ~ '
SBR4-gel 3	- Care	,	~~	****	30	-
SBR3-gel				***		30
Carbon black	70	70	70	70	70	70
Zinc white	3		3	3	3.	3
Stravio acid	1	2	3	3 3	1	3
Antioxidan:	.3.	3	3.	1	1	3
Aromatic of	35	15	25	ß	3.5	15
Saltur	1.75	1,75	1.75	1,75	1.75	1.7 5 1
Accelerator	1	1	1	1	3	1
tend (O° C.)	0.450	0.457	0.454	0.509	0,513	0,51
tand (80° C)	0,272	0.333	0.325	0.265	0,250	0.37
tsකර පන්නෙහැ	3,658	2.377	1,397	2,923	1.975	1.86

7

TABLE II continued

	Standard Ex. 2	Comp. Ex. 3	Comp. Ex. 3	Ex. 3	Ex. 4	Ka: 5
Abrasios resistance (index)	100	82	84	. 93	98	99

Standard Examples 3 to 5, Examples 6 to 7, and Comparative Examples 4 to 5

Rubber compositions having the formulations (parts by weight) shown in Table III were obtained by blending in two steps, i.e., the first step and the second step, by analogous methods to Example 1 and Comparative Examples 1 to 2 using the rubber components shown in Table V and the physical properties were evaluated. Note that the blending of the first step consisted of mixing and kneading by analogous methods to Example 1 and Comparative Examples 1 and 2, while the blending of the second step consisted of mixing and kneading the compounding agents other than the vulcanization system by a Banbury mixer, discharging the resultant mixture, then adding the vulcanization compounding agents and mixing and kneading the same. The results are shown in Table

SBR3 and SBR4: As explained above.

58R5: Styrene-butadiene rubber (Japan Elastomer, Assprene 303, Tg. ~33° C.)

SBR4-gel 1 and SBR4-gel 2: As explained above.

SBR3-gel 1: A gelled rubber prepared from styrene-butadiene rubber (Japan Elastomer, Asaprene 303) in an analogous method to SBR1-gel 1. The swelling index

SBR5-gel 2: A gelied nubber prepared from styrenebutadiene nubber (Japan Elasiomer, Asaprene 303) in an analogous method to SBR2-gel. The swelling index was 17.

8

Standard Example 7 and Examples 8 to 13

Rubber compositions were obtained by blending by analogous methods to Example I, except for using the formulations (parts by weight) shown in Table IV and evaluated the physical properties. The results are shown in Table IV.

SBR3 and SBR4: As explained above.

SBR4-gel 3: A gelled rubber prepared by adding to 400 g
of styrene butadiene rubber (Nippon zeon, Nipo) 9529, Tg:
-21° C.) 10 g of 1,10-decapethiol and mixing the resultant
mixture by a kneader at 185° C. for 40 minutes. The swelling
index was 21.

RBR4-gel 4: A gelled rubber prepared by adding to 400 g
25 of styrene-butadiene rubber (Nippon Zeon, Nipol 9529, Tg:
—21° C.) 9 g of trithincyanulic acid and mixing the resultant
mixture by a kneader at 190° C. for 15 minutes. The swelling
index was 32.

SBR4-gel 5: A gelled rubber prepared by adding to 400 g of styrene-butadiene rubber (Nippon Zeon, Nipol 9529, Tg. -21° C.) 7 g of dicumyl peroxide and mixing the resultant mixture by a kneader at 175° C. for 5 minutes. The swelling index was 15.

SBR4-gel 6: A gelled nubber prepared by adding to 400 g of styrene butadiene rubber (Nippon Zeon, Nipol 9529, Tg: -21° C.) 7 g of benzoquinone and mixing the resultant mixture by a kneader at 180° C. for 20 minutes. The swelling index was 18.

SBR4-get 7: A gelled rubber prepared by adding to 400 g of styrene butadiene rubber (Nippon Zeon, Nipol 9529, Tg: -21° C.) 11 g of phenol and 10 g of maleic anhydride and mixing the resultant mixture by a kneeder at 180° C. for 30 minutes. The swelling index was 17.

SBR4-gel 8: A gelled number prepared by adding to 400 g of styrene butathene rubber (Nippon Zece, Nipol 9529, Tg: -21° C.) 11 g of hydroquinous and 10 g of maleic anhydride and mixing the resultant mixture by a knowler at 180° C. for 40 minutes. The swelling index was 29.

TABLE III

	Skaadari Ex. 3	Standard Ex. 4	Comp. Ex. 4	Ex. s	Standard Ex. 5	Standard Ex. 5	Comp. Ex. 5	£z, 7
lst step	CONTRACTOR OF THE STREET							
SERS	70	70	סל	70	30	70	73	70)
SBR4	45	,		~~~	(444)	***		;
535.85					30	****		
Carpos black	70	70	70	70	70	70	.50	70
Zina white	3	3	.3	3 3	3 3	3	3	3
Steanio acid	1	1	1	2		2	1	3 1 1
Antioxidant 60	Ţ.	ĩ	1	Ä,	3	3	1	
Aromanic all	Q.	ខ័	15	15	3.5	15	3.5	3.5
NP	160	260	1,60	160	160	380	160	1,60
2nd step			•					
SBRA		45	•••			AMERICA.	****	inn
SBR4-Gel 1	***	****	30	****	•••		***	***
SBR4-Get 2	u.	••••	*	30	~~	,	~	-
\$BR\$	***	-	Service 1	A-1-1-1	cm.	30	****	Search
SBR5-Gel 1		and a	'www	****	, error		30	
\$19 K5-Get 2		-	****	****				30
Sulfa	1.75	3.78	3.75	1.75	1.75	1,75	1,75	1.75
Accelerator	3	3	1	Ĺ	1.	3	1	i
සාර් (0° C.)	0.450	0.415	0,460	0,475	0.445	8,418	0.450	0.468
208 (60° C.)	0.272	0.219	0.295	0.223	9,919	6.393	0.333	0,239
tand balance	2.654	1.895	2.539	2.149	1.398	1.647	1.373	1.958
Abrasion resistance (index)	100	102	83	97	100	136	86	125

ŷ

TABLE IV

COCCO, Actor and Coccoo and Cocco	Spendard		•••••	••••••	*******	***************************************	•••••••••••••••••••••••••••••••••••••••
	£x. 7	Bx. 8	Ex. 9	Ex. 10	Bx. 11	5x. 13	Rx. 13
lat step		.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,				At Maria	
SBR3	70	70	70	70	70	70	79
SBR4	25					•	
SBR#-gel 3	***	30	,	***			
SBR4-gel 4	0000		30		***		
SBR4-grl S		•••	•	30	****	***	****
SBR4-ge) 6					30	****	Admir.
SBR4-gel 7					CONT.	33	·····
8 123-4KB2	****	1	1000	· .			30
Carbon black	70	70	75	70	76	70	70
Zine white	3	3	3	3	3	3	3.
Steamin and	1	3	1	1	1	3	1
Assignifient 6C	1	3	3	3:	2	1	ž
Aromenic off	~~~	3.5	15	15	23	2.5	15
Suliu	1.75	1.75	1.75	3.75	1.75	1.75	1.75
Vulcanization accelerator	3	3	1	1	3	3 .	1
tanó (0° C.)	0.466	0,452	0.455	5.515	0.505	0.533	0,535
und (60° C.)	0.296	0.268	0.279	0.293	8,273	0.272	0.287
tanà balance	1,574	3.687	3.531	3,7793	1.863	1.900	1.864
Abrasion resistance (index)	100	28	95	87	86	87	36

ingredients in two steps, the first step and second step, by analogous methods to Examples 6 to 7, except for using the

Formulations (parts by weight) shown in Table V and evaluated the physical properties. The results are shown in Table V.

gradients in two steps, the first stem and executed the physical properties. The results are shown in Table V.

10

SBR4-gel 3 to SBR4-gel 8: As explained above.

TABLE V

	Standard Ex. 3	Steadard Ex: 9	E4. 14	Ex. 15	Ex. 16	Ex. 17	Ex. 18	Ex. 19
3st step	***************************************						************	**********
SER3	70.	70	70	70	70	70.	70	70
SBR4	45				m	•••		-
SERS			vine.	٠	****			
Carbon black	70	73	70	79	70	70	38	70
Zine white	3	3	3	.3	3	3	3	3
Steams sold	3.	1	1	ž	1	1	3	3
Asticatidant &C	1	1	1	1	-2	3.	ž	1
Aromskie od	3	0	15	2.5	3.5	1.5	15	7.8
NE	160	360	160	160	160	150	3.80	1.89
Ind step								
SER4		45	****	***			****	****
5884-04 3			30				***	
SBR4-Ge) 4	***	***		30		****	****	12000
SBR4-Gel S	***	****		***	30			***
SBR4-Gel 6	r000'					30	•••	
\$BR4-Gel 7				-	,	•	30	
SBR4-Gel 8		***	·					30
Salfue	1.75	2.75	1.75	1.75	3.75	1.75	1.75	1.75
Vulcanization sconfermor	1	2.	i	1	1	1	1	1
tend (0° C)	0,466	0.414	0.443	0.419	0.513	0.510	6.507	0.485
taző (60° C.)	0.296	0.234	0.233	0.233	0.285	0.246	0.241	0.241
tanà balance	1,574	1.769	1,883	1,798	1,800	2.072	2,103	2.013
Abresios resistante (iudex)	100	101	92	110	102	308	137	109

resultant mixture.

11

As explained above, the rubber composition containing the gelled rubber according to the present invention can provide a rubber composition having a high tan 8 at 0° C. (i.e., good wet brake performance) and a low tag & at 60° C. (i.e., good rolling resistance) which is suitable for applica- 5 tions such as fires as a subber composition superior in viscoelasticity. Further, it can be used for conveyor belts, rubber hoses, fenders, rubber sheets etc.

What is claimed is:

1. A subper composition comprising 50 to 90 parts by 10 weight of a diene rubber and 50 to 10 parts by weight of a galled rubber having a toluene swelling index of 15 to 150, based upon the total amount of 100 parts by weight of the diene rubber and the gelled rubber, wherein a glass transition temperature of the diene rubber is at least 10°C, lower than 15 gelied rubber is 16 to 150. a glass transition temperature of the gelled rubber.

2. A rubber composition as claimed in claim 1, the gelled rubber contains at least 0.1% by weight of an acid anhydride

molety.

3. A process for producing a rubber composition comprising 100 parts by weight of a starting rubber containing 30 to 90 parts by weight of a diene rubber and 50 to 10 parts by weight of a golled rubber, the glass transition temperature of the diene rubber being at least 10° C. less than the glass transition temperature of the gelled rubber, and 30 to 120 parts by weight of a filler, comprising the steps of mixing the diene rubber and at least 80% by weight of the total amount of the filter at a temperature of at least 135° C., and then mixing the galled rubber and the remaining filler with the

12

4. A process for producing a rubber composition as claimed in claim 3, wherein the toluene swelling index of the